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Efficiency enhancement and doping type inversion in Cu_2CdSnS_4 solar cells by Ag substitution⁺

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The main limiting factor of kesterite-related solar cells is the low opencircuit voltage (V_{OC}) relative to their bandgap. This drawback has been correlated with the easily formed anti-site defects caused by similar ionic radii and/or chemical valence. Recent success in suppressing Snrelated defects by using Cd to replace Zn in Cu₂CdSnS₄ was attributed to the higher formation energy of the $2Cu_{Cd} + Sn_{Cd}$ defect complex compared to its counterpart in Cu₂ZnSnS₄. This has motivated the use of bigger ions to replace Cu in Cu₂CdSnS₄ to reduce the possibilities of I_{II} and II₁ defect formation. In this work, we substitute Cu in Cu₂CdSnS₄ with larger Ag at various concentrations and investigate the structural, optoelectronic, and photovoltaic properties of (Cu,Ag)₂CdSnS₄. Higher concentrations of Ag lead to peak splitting in XRD spectra, which is attributed to mixed phases and marks the transition towards Ag₂CdSnS₄. This is also complemented by Raman scattering analysis, the first time the Raman spectrum of Ag₂CdSnS₄ is reported. Doping type inversion was observed for pure n-type Ag2CdSnS4 instead of the p-type of Cu₂CdSnS₄, accompanied by high carrier mobility and sharp absorption onset. Further optoelectronic and photovoltaic characterization reveals that adding 5% Ag concentration improves Cu₂CdSnS₄ device performance to 7.72%, mainly due to superior film quality and improved interface properties. As a result, better carrier collection contributes to the short-circuit current improvement of the champion device.

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Introduction

The motivation to address anti-site defects in Cu₂ZnSnS₄ has led to investigations on I2-II-IV-VI4 compounds as better solar harvesting materials.^{1,2} Anti-site defects in Cu₂ZnSnS₄ occur when one cation replaces another with similar or different oxidation states, resulting in point defects such as Cuzn, or defect clusters such as 2Cu_{Zn} + Sn_{Zn}. To mitigate the formation of these defects, cations with larger ionic size differences are utilized to substitute one or more cations with similar ionic sizes, such as Cu or Zn.3

Cation substitution in Cu₂ZnSnS₄ is a crucial strategy in further improving its performance. The cation substitution on Cu_2ZnSnS_4 involving the Ag & Cd reduces the V_{OC} deficit caused by both radiative and nonradiative recombination. Partial Cdalloying of Cu₂ZnSnS₄ has positively impacted grain quality,⁴ band alignment,⁵ and bandgap fluctuations.⁶ As indicated by lower bandgap fluctuation, Cd substitution limits the subbandgap absorption compared to Cu₂ZnSnS₄ and elevates the valence band maximum (VBM), improving the band alignment.6,7 Similarly, Ag substitution has a positive effect, as indicated by lower Urbach energy and reduced anti-site defects.8 Furthermore, the co-substitution of Ag and Cd on Cu₂ZnSnS₄ improved device efficiency to 10.2%.9

Full cation substitution of Cu₂ZnSnS₄ leading to I₂-II-IV-VI₄ compounds has also been explored to pursue different structural and electronic properties. The lack of Cu vacancy in Ag₂-ZnSn(S,Se)₄ and Zn-rich condition was found to be favorable for forming n-type instead of p-type material.¹⁰ Consequently, a different device configuration needs to be employed.11,12 CXTS compounds (where X = Mg, Mn, Ni, Co, Fe, *etc.*) have been studied as substitutes for Zn. A library of CXTS compounds was successfully fabricated using spray pyrolysis. While Cu₂MnSnS₄, Cu₂BaSnS₄, and Cu₂SrSnS₄ demonstrated good device performances, forming pure phases for some compounds was difficult.13 Efforts have also been made to alloy a high amount of Ge for CZGSe, leading to a 7.6% efficiency and a further 8.5% after surface treatment and heat treatment on a complete device

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(including MgF_2 anti-reflection coating layer).^{14,15} Among all these cations, Cu_2CdSnS_4 has the highest device performance so far.

The favorable material properties of Cu₂CdSnS₄ make it a promising candidate for further development as one of the emerging thin film solar cells. The Cu₂CdSnS₄ solar cell has shown remarkable device performance, starting with an efficiency of 7.96%, which recently has been improved to 10.1%, placing it among the highest reported efficiency of emerging kesterite-inspired solar cell devices and close to the 11.0% certified record efficiency for pure sulfide Cu2ZnSnS4.16-19 The suppressed 2Cu_{Cd} + Sn_{Cd} defect formation, due to its higher formation energy compared to $2Cu_{Zn} + Sn_{Zn}$ in Cu_2ZnSnS_4 , is attributed to the promising properties of this material. Both defect complexes act as deep electron traps3,20 and contribute to the increase of bandgap fluctuation and nonradiative recombination. Cu-Poor condition, commonly employed in kesterite solar cells, further suppressed the 2Cu_{Cd} + Sn_{Cd} defect.¹⁶ This non-stoichiometric condition and optimized sulfurization configuration were also crucial for the recent device efficiency improvement.17 While the current achievements for Cu2CdSnS4 solar cells are encouraging, additional strategies such as doping and alloying could be important to understand this compound better.

The cation substitution (and doping) strategies in Cu₂-CdSnS₄ have yet to be thoroughly studied. While the beneficial roles of Ag have been observed in partially Cd-substituted Cu₂ZnSnS₄ compounds, its impact on the material and photovoltaic properties of Cu2CdSnS4 remains unexplored.9,21 Additionally, the absorption coefficient of Ag₂CdSnS₄ has been calculated to be suitable for a solar cell.²² The larger Ag⁺ ionic radii (1.00 Å) compared to Cu⁺ (0.60 Å) suggest that substituting Cu to Ag could further reduce $2Cu_{Cd} + Sn_{Cd}$ defects. Other theoretical works highlighted that as the difference in ionic radii size of Ag–Cd is not as high as Ag–Zn (Ag⁺ to Cd²⁺ is 28% larger, while Ag⁺ to Zn²⁺ is 66% larger), Ag₂CdSnS₄ might not be as disorder-resistant as compared to Ag₂ZnSnS₄.^{23,24} This work aims to investigate the effect of Ag substitution for Cu₂CdSnS₄ by comparing phase distribution, film morphology, and device performance as part of its structural, optoelectronic, and photovoltaic properties. The small addition of Ag in Cu₂CdSnS₄ improved device performance to 7.72% compared to 6.87% reference, but the improvement is limited by the presence of mixed phases at higher Ag concentrations. This limitation will be explored by studying the transformation from Cu₂CdSnS₄ to Ag₂CdSnS₄, correlating the transition with the evolution of the device performance. Finally, the fully-substituted Ag₂CdSnS₄ induced doping type inversion, high carrier mobility, and exhibits a sharp absorption onset.

Results and discussion

In this study, we monotonically modify the concentration of Ag by depositing the precursor solutions using spin-coating, starting from pure Cu_2CdSnS_4 (Ag/[Ag + Cu] = 0%) to Ag_2-CdSnS_4 (Ag/[Ag + Cu] = 100%). The selection of 5% and 10% Ag was based on the optimal device efficiency observed in previous

 Cu_2ZnSnS_4 devices.^{25–28} The films for all Ag concentrations were characterized for their structural and optoelectronic properties, while only selected concentrations were made into full devices.

Structural properties of (Cu_{1-x}Ag_x)₂CdSnS₄

The diffractograms (Fig. 1a) obtained from as-fabricated Cu₂-CdSnS₄ film fit the reported database and showed major peaks at 28° and 47° associated with (112) and (204) of the Cu₂CdSnS₄ stannite plane (JCPDS No. 29-0537). The magnified XRD spectra in the range of $2\theta = 23-30^{\circ}$ is shown in Fig. 1b, highlighting the transition from tetragonal Cu₂CdSnS₄ to orthorhombic Ag₂-CdSnS₄ phase. In the range of 0–20% Ag substitution, there is a subtle shift to a lower diffraction angle at the (112) peak due to the successful incorporation of larger Ag ions.²⁹ The slight 40° diffraction angle is from the Mo substrate and thus is used as a reference for the peak shifting for other Ag concentrations. It is observed that the transition from Cu₂CdSnS₄ towards mixed phases with Ag₂CdSnS₄ starts at 30% Ag substitution. Starting at this concentration up to 80% Ag, it can also be observed that the film quality is deteriorating as the main peak intensity is relatively low compared to the Mo peak as substrate. The shifting trend of the (112) peak of Cu₂CdSnS₄ continued to higher Ag concentrations too. The primary peaks at 25.06°, 26.54°, and 28.48° associated with (011), (200), and (111) of the Ag₂CdSnS₄ planes are observed, respectively.

To identify the presence of secondary phases, Raman measurement for all films were conducted as shown in Fig. 1c. Under 532 nm excitation wavelength, the Cu2CdSnS4 film exhibits its characteristics symmetry modes at 332 and 283 cm⁻¹, similar to previously reported Raman data.16,17 The appearance of these two peaks dominates up to 20% Ag. Note that the CdS secondary phase can also be identified in the spectra, especially at the 30-80% Ag films, where the film quality seems to deteriorate. On the other hand, no Cu₂SnS₃ secondary phase (290 and 350 cm⁻¹) is detected on all films. At the higher amount of Ag (\geq 60%), the main peak at 332 cm⁻¹ shifts to a higher wavenumber value, and the appearance of a shoulder peak at 367 cm⁻¹ suggests a change in the phase from the stannite Cu₂CdSnS₄ to orthorhombic Ag₂CdSnS₄. Although to our knowledge Raman studies about Ag₂CdSnS₄ have not been reported, K. Pietak et al. presented in their study of Ag₂ZnSnS₄ similar results regarding their Raman measurements with the appearance of an intense peak at 341 cm⁻¹ as well as an additional peak at 370 cm^{-1} .³⁰ In addition to our XRD, it seems that the observed Raman peaks phase belongs to orthorhombic Ag₂CdSnS₄, the first time it has been reported in the literature.

The transition between Cu_2CdSnS_4 into Ag_2CdSnS_4 was also quantitively studied by Rietveld refinement of the XRD spectra. All films have been refined to their tetragonal unit cells for Cu_2CdSnS_4 , orthorhombic for Ag_2CdSnS_4 , and both for the partial Cu–Ag substitution. The value of R_{bragg} evaluates how well each phase fits with the model, the lower, the better. The goodness of fit (GOF), representing the closeness of the refinement model to the actual measured data with one as the ideal value, is used to determine the refinement model quality. The GOF equation is shown below,

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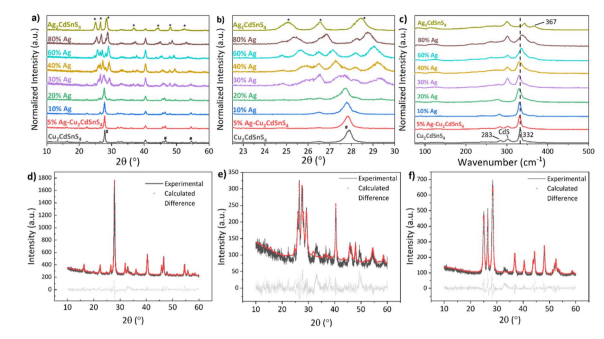


Fig. 1 (a) X-ray diffraction (XRD) pattern and (c) Raman spectra of Cu_2CdSnS_4 film with various Ag content and Ag_2CdSnS_4 film. (b) Magnified XRD view on 23–30° diffraction angle near (112) peak of Cu_2CdSnS_4 ; the intensities in (a–c) are normalized, % Ag refers to the concentration ratio of Ag/Ag + Cu. (d) Refinement plot for Cu_2CdSnS_4 sample, (e) 30% Ag of $(Cu_{1-x}Ag_x)_2CdSnS_4$ sample, and (f) Ag_2CdSnS_4 sample.

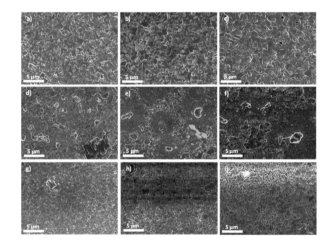


Fig. 2 Surface SEM images for (a) Cu_2CdSnS_4 and additional Ag content of (b) 5%, (c) 10%, (d) 20%, (e) 30%, (f) 40%, (g) 60%, (h) 80%, and for (i) Ag_2CdSnS_4 .

$$\text{GOF} = \chi^2 = \left[\frac{R_{\text{wp}}}{R_{\text{exp}}}\right]^2$$

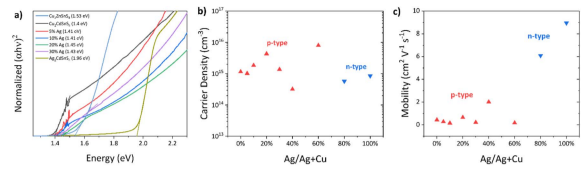
where R_{wp} is the residual weighing the fitting of the model to the peak intensities, and R_{exp} is the residual for the experimental value evaluation.³¹

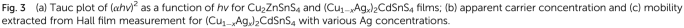
The refinement for pure Cu₂CdSnS₄ sample can be closely fitted ($R_{\text{bragg}} = 3.79$; GOF = 1.53) to the calculated tetragonal unit cells, as shown in Fig. 1d. This fitting case still applies after adding 5% up to 20% Ag (Fig. S1†). At 30% Ag, the experimental data shown in Fig. 1e cannot be reasonably fitted to tetragonal

Cu₂CdSnS₄ ($R_{\text{bragg}} = 5.27$) or orthorhombic Ag₂CdSnS₄ ($R_{\text{bragg}} = 9.66$). Finally, the 80% Ag ($R_{\text{bragg}} = 2.72$; GOF = 1.47) and pure Ag₂CdSnS₄ ($R_{\text{bragg}} = 2.95$; GOF = 1.46) data have a close fit to the calculated orthorhombic unit cells, as shown in Fig. 1f and S1.†

Following, the study on film morphology (Fig. 2) is carried out. Pure Cu₂CdSnS₄ and the Ag-alloyed films have large and compact grains with up to a 10% Ag concentration ratio. Meanwhile, the grains become smaller and less uniform between 20% and 40% Ag. While the presence of grain boundaries could be benign for high efficiency Cu(In,Ga)Se₂ thin film solar cells due to the effective passivation by alkali,^{32,33} in general for polycrystalline material, such as the more closely related Cu₂ZnSnS₄, grain boundaries may introduce additional deep level states within the energy bandgap acting as effective recombination centre.^{34,35} The grains become more uniform as the Ag content increases to 60%. Finally, the Ag₂CdSnS₄ film has uniform grains, albeit smaller than the Cu₂CdSnS₄ film. This means the uniform grains that could benefit a solar cell device can be obtained from pure Cu₂CdSnS₄ and Ag₂CdSnS₄ film, as well as the 5% Ag.

The elemental ratios of the films were measured by EDS, as shown in Table S1.[†] All the films have non-stoichiometric Cupoor and Cd-rich compositions similar to the as-prepared solutions. The Ag concentration in each variation is in good agreement with the prepared amount. A line scan was conducted for 5% and 30% Ag films to check the elemental distribution of the film. Ag is distributed uniformly throughout both films, similar to other cations, as shown in Fig. S2.[†] Thus, there is no Ag distribution grading for both the small and mixed phases concentrations.





Optoelectronic properties of (Cu_{1-x}Ag_x)₂CdSnS₄

UV-Vis was conducted for all concentrations to understand further the optoelectronic properties of the $(Cu_{1-x}Ag_x)_2CdSnS_4$ films. The increased bandgap for $(Cu_{1-x}Ag_x)_2CdSnS_4$ in Fig. 3a and S3[†] can be attributed to the higher incorporation of Ag, as demonstrated by the peaks shifting observed from XRD. The absorption edge for pure Cu_2CdSnS_4 and small concentrations of Ag up to 10% are quite sharp, indicating their better light absorption ability, while it is less so for 20% Ag and 30% Ag. At 30% Ag, the mixed phases clearly contributed to the less sharp absorption edge. This observation is the same for higher Ag concentrations, but for pure Ag_2CdSnS_4 and Cu_2ZnSnS_4 can be obtained with an extracted bandgap of 1.93 eV.

Hall measurement is then conducted for all concentrations of the $(Cu_{1-x}Ag_x)_2CdSnS_4$ films. The apparent carrier concentration for Cu_2CdSnS_4 is 2.7×10^{15} cm⁻³ with carrier mobility of 0.44 cm² V⁻¹ s⁻¹. The presence of mixed phases from 30% to 80% Ag could affect the values of their carrier density, as there is no clear correlation between carrier concentration and mobility values to the Ag concentrations, as observed in Fig. 3. Interestingly, the doping type changes between p-type for Cu_2CdSnS_4 into n-type for Ag₂CdSnS₄. The high formation energy of its acceptor defects could be the reason for this transition, as similarly observed in Ag₂ZnSnS₄ and Ag₂ZnSnSe₄.^{11,12}

The highest peaks obtained from photoluminescence spectra in Fig. S4a[†] do not represent the bandgaps of all the samples. However, secondary peaks around 1.35 eV close to each respective bandgap value of 5% and 10% Ag films were observed, indicating fewer defects or improved quality after small doping of Ag. The main peaks for higher concentrations of Ag (30%, 60%, and pure Ag_2CdSnS_4) were all blue-shifted corresponding to their higher bandgaps.

Photovoltaic properties of (Cu_{1-x}Ag_x)₂CdSnS₄

The complete device configuration is SLG/Mo/ $(Cu_{1-x}Ag_x)_2$ -CdSnS₄/CdS/ITO without anti-reflection coating. *I–V* measurements were conducted on at least ten devices for each Ag concentration to obtain statistical representation. The photovoltaic parameters for the devices up to 60% Ag are presented in Fig. 4. Between 0% to 10% Ag, there is no significant improvement in FF and V_{OC} . The performance starts to drop at 20% Ag concentration, accompanied by a drop in J_{SC} and FF, which can be related to the presence of non-uniform grains and voids on the film. The presence of mixed phases between 30% and 40% Ag is well-reflected in the device performance. While for 30% Ag, the V_{OC} is comparable with lesser Ag concentration, at 40%, the V_{OC} drops significantly.

Both 30% and 40% of Ag devices have poor FF and J_{SC} . The presence of pinholes might be responsible for the poor V_{OC} and FF at those concentrations. At the same time, the J_{SC} also suffers from carrier recombination at the grain boundaries due to poor grain growth on top of the increasing bandgap. The devices have very poor performance when Ag content is higher than Cu at 60%, which could be attributed to the much higher bandgap affecting the band alignment with the CdS layer. Moreover, despite improved morphology conditions at 80% and 100% Ag, an alternative device configuration might be necessary to conform to the n-type majority carriers' condition, as observed in Fig. 3. Lastly, we did not obtain a working Ag₂CdSnS₄ device because the remaining pure phase is n-type at this concentration.

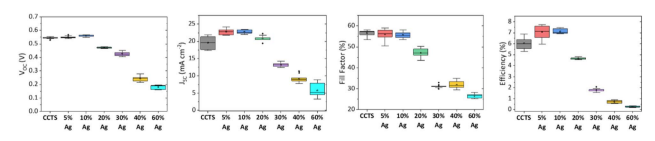


Fig. 4 Device parameters for Cu₂CdSnS₄ with various Ag concentrations. All boxplots represent ten devices each.

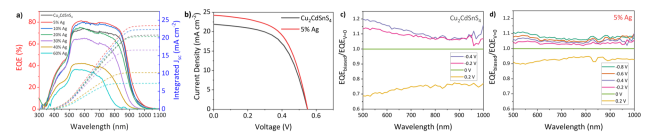


Fig. 5 (a) External quantum efficiency (EQE) curves (left axis) and their integrated J_{SC} (right axis-dashes) for Cu₂CdSnS₄ with various Ag concentrations, (b) I-V curves for the best Cu₂CdSnS₄ and 5% Ag device. Voltage-biased EQE curves for (c) Cu₂CdSnS₄ and (d) 5% Ag device normalized to the unbiased EQE values.

Table 1	Photovoltaic parameters of the	e best Cu ₂ CdSnS ₄ and 5% Ag device
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Device	$V_{\rm OC} [{\rm mV}]$	$J_{\rm SC} \left[{ m mA~cm^{-2}} ight]$	FF [%]	η [%]	$E_{\rm g} \left[{\rm eV} ight]$	п	$J_{01} \left[\times 10^{-6} \mathrm{A} \mathrm{cm}^{-2} ight]$	$R_{\rm s} \left[\Omega \ {\rm cm}^2 \right]$	$G_{ m sh} \left[m mS \ m cm^{-2} ight]$
Cu ₂ CdSnS ₄	549	21.89	57.16	6.87	1.39	2.35	2.23	2.55	3.93
5% Ag	550	24.23	57.89	7.72	1.41	2.64	6.64	1.63	3.79

The External Quantum Efficiency (EQE) for the bestperforming devices up to 60% Ag concentration is shown in Fig. 5a. Poor EQE results were obtained for Ag concentrations above 60%. The J_{SC} improvement for 5% and 10% Ag devices is well-reflected by each device's higher EQE values, which is translated to the higher integrated J_{SC} values. For 5% Ag, the EQE increases throughout the whole wavelength. This improvement can be attributed to the reduced recombination at the bulk of Cu₂CdSnS₄ due to small Ag content doping.³⁶ Similar improvement is observed from the 10% Ag, but there is a reduction at the lower wavelength, probably due to parasitic absorption of the top layers from CdS/ITO.37 The higher Ag contents are consistent between J_{SC} values and the EQE. At higher concentrations, the bandgap increases to 1.63 eV for 60% Ag. This increasing trend is also obtained from the extracted bandgap values from the absorption spectra in Fig. S3.[†] Moreover, this is also consistent with the experimentally observed 1.93 eV bandgap for pure Ag₂CdSnS₄.³⁸

The improvement for the best device with 5% Ag mainly comes from slightly higher J_{SC} and Fill Factor (FF) compared to the pure Cu₂CdSnS₄ devices, as observed in Table 1 and Fig. 5b. The better FF for the 5% Ag device is in good agreement with its lower R_s and G_{sh} value. The same J_{SC} improvement for the small amount of Ag was also observed in CZTS devices with a high Cd substitution concentration. It was attributed to larger grain sizes and enhanced depletion width.⁹

To further understand the reason for the improved performance of the 5% Ag device, we performed a C-V measurement. Comparing the pure Cu₂CdSnS₄ and 5% Ag devices, as shown in Fig. S5,† the extracted carrier concentration for these devices does not significantly change, as similarly observed from Hall measurement (Table S2†). The amount of Ag might be too small to significantly affect the carrier density. Nevertheless, the 5% Ag device shows a slight improvement in its depletion width, leading to a better carrier collection, as reflected by its superior EQE. We also performed EQE measurements under positive and negative voltage bias for the two devices, as shown in Fig. 5c and d. The EQE_{biased}/EQE_{V=0} ratio indicates severe interface recombination in Cu₂CdSnS₄. In particular, the large deviation of this ratio from the ideal value of 1 and the corresponding slope of the dependence of this ratio on wavelength suggests the presence of interface recombination.³⁶ Adding 5% Ag to Cu₂-CdSnS₄ brings this ratio closer to 1, suggesting that interface recombination in these devices is suppressed to some extent. The 5% Ag device performance improvement can be mainly attributed to this reason and the better film quality as demonstrated by the presence of a PL peak at the energy level close to its bandgap value.

The EQE spectrum onset for Cu₂CdSnS₄ and 5% Ag devices EQE))/dE as presented in Fig. S6[†] referring to previous methods in the literature.^{39,40} According to Mattheis et al.,³⁹ the standard EQE))/dE represents the amplitude of bandgap fluctuations $(\sigma_{E_{\sigma}})$. For the 5% Ag device, the $\sigma_{E_{\sigma}}$ value of 56 meV is not far from the σ_{E_a} value of the Cu₂CdSnS₄ device (58 meV). However, these values are much smaller than our similarly processed Cu₂ZnSnS₄ solar cell device (116 meV) and comparable to the σ_{E_a} value of the current 13.8% record efficiency Cu₂ZnSn(S,Se)₄ (CZTSSe) solar cell device, which is about 52 meV.41 As the degree of bandgap fluctuations has been associated with the $2Cu_{Cd} + Sn_{Cd}$ (or $2Cu_{Zn} + Sn_{Zn}$) deep defects, ^{3,16,42} the small $\sigma_{E_{a}}$ values of Cu₂CdSnS₄ and 5% Ag devices indicate the suppression of the associated deep defects. A combination of this work and the use of Sn⁴⁺ precursors (used in the 13.8% CZTSSe device and currently regarded as essential for high-efficiency CZTSSe solar cells) might give better CCTS device performance in future studies.

Conclusions

In this work, we have investigated Ag substitution on Cu_2 -CdSnS₄ and monotonically studied its transition. The phase transition from tetragonal Cu_2CdSnS_4 to orthorhombic Ag₂CdSnS₄ starts at 30% Ag concentration, as observed from the XRD spectra. Despite the bigger ionic radii size (Ag 1.00 Å vs. Cu 0.60 Å, Cd 0.78 Å), higher Ag concentrations do not effectively translate into better photovoltaic performance. This is demonstrated by its poor crystal quality, optoelectronic properties, and possible carrier-type changes. However, the n-type nature of the fully-substituted Ag₂CdSnS₄ might require a different device configuration and further optimization to obtain a working solar cell device. On the other hand, small amounts of Ag concentrations of up to 10% have improved device performance compared to pure Cu₂CdSnS₄. The 5% Ag device has the best performance of 7.72%, which is attributed to its better film quality and superior interface properties, leading to its higher *J*_{SC} compared to the pure Cu₂CdSnS₄.

Author contributions

A. I. fabricated the $(Cu_{1-x}Ag_x)_2CdSnS_4$ thin films, conducted XRD and its Rietveld analysis, carried out SEM-EDS, UV-Vis, Hall measurement and solar cell device characterization, and analyzed the raw data. S. L. helped with the analysis of solar cell device characterization and Hall measurement results. J. M. R. T. contributed suggestions on the synthesis. R. S. performed the PL measurements, EQE-V, and helped with the analysis of the results (supervised by R. A.). A. G. M. conducted Raman measurement and helped with the discussion of the result (supervised by E. S.). S. H. contributed suggestions on EQE-V data. L. H. W. supervised the work.

Conflicts of interest

There are no conflicts to declare.

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